# SPECIFICATION



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## COMPLETE SPECIFICATION

## Method for Agglomerating Vulcanization Accelerators

We, THE GOODYEAR TIRE & RUBBER COM-PANY, a corporation organized under the laws of the State of Ohio, United States of America, with offices at 1144, East Market Street, Akron, Ohio, United States of America, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed, to be particularly described in and by the follow-10 ing statement:-

This invention relates to powdered vulcanization accelerators in a new and more useful form. More specifically, it relates to methods for agglomerating certain powdered accelera-

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The use of powdered organic accelerators in the compounding of natural and synthetic rubber presents certain problems to the rubber goods manufacturer. For example, the finely divided nature of the accelerator causes it to be blown into the air during production opera-tions such as milling. This results in an increase in the cost of production and in the possible contamination of other materials. Then, too, certain of these accelerators create serious health hazards in production operations because once blown into the air surrounding production operations, they cause irritation to the skin and nasal passages of the workmen exposed to the contaminated air. In addition, the fine powder-like nature of the pigment makes uniform dispersion difficult since the pigment tends to form lumps and flakes which stick to or cake on the mill rolls, thus increasing the time and effort required to form an intimate and thorough dispersion of the pigment in the rubber.

Efforts have been made to reduce the objectionable features of the powdered accelerators by converting the finely divided pigment particles into some form of agglomerate. It has usually been found that, in order to form such agglomerates, some additive must be used with the powdered pigment if the agglomerate, once formed, is to be stable. Frequently the additives are materials which may be objectionable when ultimately mixed into the rubber compound with which the accelerator is to be used Other additives produce agglomerates which are cohesive and not free-flowing, with the result that such agglomerates do not lend themselves to normal handling and weighing procedures. Other additives employed for preparing agglomerated accelerators are used in such large amounts that the rubber chemist 55 is required to work with an excessively "diluted" accelerator. Other treatments produce agglomerated accelerators which do not disperse readily into the rubber during the milling operation. Still other treatments result 60 in an agglomerated form which is not sufficiently stable to maintain its agglomerated form while it is transported from the manufacturer to the user. All of these objections have been overcome by preparing the powdered accelerators in an agglomerated form according to the methods of this invention.

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One object of this invention is to provide for the preparation of powdered accelerators in an agglomerated form which will substantially eliminate the tendency of such accelerators to form dust which is blown into the surrounding air during storage, shipping, weighing, and processing operations. The use of these novel forms of accelerator results in a material saving to the manufacturer, the elimination of a source of contamination, and the elimination of a health hazard to the workman handling such materials.

Another object of this invention is to provide for the preparation of these accelerators in an agglomerated form of such a nature that the individual particles of the agglomerate will hold together during normal handling operations and yet will disperse uniformly into and through the rubber during the milling operation. Still another object is to provide for the preparation of agglomerated organic rubber vulcanization accelerators which disperse more rapidly into the rubber during milling than will the unagglomerated powdered accelerator additional object

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accelerators which ...prepare, agglomerated free - flowing are and not COhesive. Another object is to prepare agglomerated organic accelerators with a minimum 5 amount of diluent. Still another object is to prepare agglomerated accelerators which contain no materials deleterious to the rubber compound with which the accelerator is to be used.

The rubber accelerators to which this invention relates are tetramethyl thiuram disulfide, tetraethyl thiuram disulfide, tetramethyl thiuram mono-sulfide, zinc dimethyl dithiocarbamate, zinc diethyl dithio carbamate, zinc di-15 butyl dithiocarbamate, mercaptobenzothiazole, benzothiazyl disulfide, and the water-insoluble metallic salts of mercaptobenzothiazole such as zinc mercaptobenzothiazole and lead mercaptobenzothiazole. For certain compounding pur-20 poses two or more of these accelerators may be employed to form a mixed agglomerate.

One type of additive used in the preparation of agglomerated accelerators is natural or synthetic rubber latex. It has been observed, 25 however, that these accelerators, and particularly mercaptobenzothiazole, when mixed with latex cause coagulation of the latex before the latex and the accelerator can be thoroughly blended together. This premature coagulation 30 prevents the production of agglomerated accelerators having the desired characteristics.

In accordance with the invention, it has been found that the objects of this invention can be accomplished by mixing an aqueous slurry of the accelerator with a small amount of a rubber latex containing an antioxident and an acidstable emulsifier, coagulating the latex after it. has been thoroughly blended with the accelerator, removing the excess water, forming

40 agglomerates, and drying the agglomerates.

The rubber latices useful in the practice of this invention are natural rubber latex, polychloroprene latex, polybutadiene latex, the latices of the rubbery copolymers of butadiene and styrene and the latices of the rubbery copolymers of butadiene and acrylonitrile. Of these, the latices of the rubbery copolymers of butadiene and styrene and the latices of the rubbery copolymers of butadiene and acrylo-50 nitrile are preferred.

The term "acid-stable emulsifier" means an emulsifier which is still effective as such at a pH below 7, as distinct from materials such as ammonia and sodium chloride which have 55 been used to stabilize latex but do not prevent coagulation on acidification.

The acid-stable emulsifiers useful in the practice of this invention will generally be organic, long chain compounds and may be anionic, cationic or nonionic in nature. The emulsifiers anionic acid-stable synthetic include (1) the metal salts of alkyl aryl sulfonates, examples of which are the sodium alkyl aryl suifonates, such as sodium dodecyl naphthalene sulfonate and similar materials sold

under the trade name Nacconol NR by National Aniline, (2) the metal salts of alkyl sulfonates, examples of which are sodium myristyl sulfonates and sodium n-decyl sulfonate, (3) the metal salts of alkyl sulfates such as sodium lauryl sulfate and sodium myristyl sulfate, (4) the metal salts of sulfated and sulfonated amides and amines such as sodium alkyl phenol ethylene oxide sulfate and, (5) the metal salts of sulfated and sulfonated esters and ethers, such as the ester of oleic acid and hydroxyethane sodium sulfate.

Materials that are cationic in nature when ionized may also be useful in the practice of this invention. Some of these cationic acidstable materials are: dipolyoxyethylene alkyl tertiary amines resulting from the condensation of ethylene oxide with organic amines, where the alkyl group is derived from fatty acids containing from 12 to 18 carbon atoms and from 2 to 50 mols of ethylene oxide are used per mol of amine. Examples of such materials are available commercially under the trade name "Ethomeens" (Registered Trade Mark) from Armour and Company, Chicago, Illinois. Another type of useful cationic material is the rosin acid substituted amine salts such as dihydroabiethylamine acetate commercially available under the trade name "Rosin Amine D" from Hercules Powder Company, Wilmington, Delaware.

Nonionic wetting agents or emulsifiers that are acid-stable are also useful in the practice of this invention. One of these nonionic materials is an alkylated aryl polyether alcohol 100 such as the material sold under the trade name "Triton X-100" ("Triton" is a Registered Trade Mark) by Rohm and Haas, Philadelphia, Pennsylvania.

In the practice of this invention it is pre- 105 ferred to use an alkyl aryl sulfonate such as sodium alkyl aryl sulfonate. With some powdered accelerators it is helpful to use, in conjunction with the acid-stable anionic sodium alkyl aryl sulfonate, a small amount of an acidstable nonionic material such as an alkylated aryl polyether alcohol or a cationic material such as a dipolyoxyethylene alkyl tertiary

In addition to the acid-stable emulsifier, the 115 latex should contain an antioxidant such as phenyl beta naphthylamine and preferably a non-discoloring antioxidant, such as the styrenated and alkylated phenols and phenyl phosphites, in order to preserve the rubber 120 content of the agglomerate against aging during storage.

The presence in the agglomerate of only a minimum amount of rubber hydrocarbon from the latex is preferred for the reason that the 125 users of the accelerator desire one which contains the least possible amount of non-accelerator material. It has been found that the latex required to produce satisfactory agglomerates of the specific accelerators mentioned above 130

should provide at least 4 parts by weight of size and shape of the particular accelerator. In rubber hydrocarbon per 100 parts by weight general, it has been found that the larger the of the powdered accelerator. While satisfactory particle size, the smaller the amount of water agglomerates can be produced using as much as 10 parts by weight of rubber hydrocarbon per 100 parts by weight of accelerator, it is preferred that from 4 to 6 parts by weight of the parts by weight of the per 100 parts by weight of accelerator, it is preferred that from 4 to 6 parts by weight of the total mix be used to minimize the dilution of the accelerator by the rubber hydrocarbon.

10 The chemical preparation of the accelerators themselves usually results in an aqueous slurry of the powdered material which can be employed in the preparation of the agglomerated accelerators without the requirement of first drying the accelerators and subsequently re-wetting them to form the slurry required for the agglomerating process. It is, therefore, economical and efficient to employ the aqueous slurry of the accelerators before the accelerator itself is dried to a powder, rather than to dry the accelerator and subsequently form the aqueous slurry. If, however, dried accelerators are employed in the preparation of the agglomerated material, it has been found to be 25 advisable to employ in the aqueous slurry a werting agent, such as the emulsifiers described above. It is also sometimes advisable to use a water-soluble salt such as sodium chloride or potassium chloride in the slurry of accelera-30 tor to aid in preventing premature coagulation

of the latex to be added later.

The latex containing the antioxidant and the acid-stable synthetic emulsifier is mixed with the aqueous slurry of the accelerator. Although 35 it is possible to use a concentrated fatex in the mixing operation, it has been found to be advisable, in order to ensure complete mixing before coagulation, to add the latex in the form of a dilute solution such as one edutating of a latex prepared from 67 parts of buradiene approximately from 5 to 10% rubber hydrocar—and 33 parts of a cylonianie using 5 parts of bon by weight. After the latex and aqueous a sodium likyl arry sulfonate as the emulsifier. slumy of accelerator have been thoroughly blended, the latex is coagulated by the addition of a small amount of a salt which will 45 either create an acid condition in the solution or will form an insoluble salt with the synthetic emulsifier. Materials which have been found to be effective for the coagulation of the latexit include alum and magnesium sulfate. Heat is 50 also helpful in the coagulation of the latex. cire

After the latex has been coagulated, the excess water is removed by decantation or filtration, tising additional water if necessary to remove the water-soluble salts from the aqueous slurry. Sufficient water is removed excess water being decanted after each washfrom the slurry to provide a mixture having a sing. After the last washing, the mixture was 120 paste-like occasistency which lends itself readily to the formation of agglomerates by means of any conventional type of pellet-form—mextruded into string-like or rod-like shapes and ing apparatus, such as a granulator, molding dried at 125°F. press, corrugated rolls or an extruder. The amount of water remaining in the paste-like mixture will depend upon the particular accelerator being agglomerated and is believed of a latex prepared from 75 parts of butadiene to be at feast in part controlled by the particle and 25 parts of styrene, using 5 parts of a

is required to provide the desired paste-like consistency, with the exact amount being determined by the particle size of the accelerator.

The preferred method for forming agglomerates is to extrude the paste-like mixture through a die provided with a plurality of small cylindrical holes. The mixture leaves the extruder in the form of springs or rods which are collected, preferably on a moving belt and dried. Excessively high temperatures should be avoided to prevent discloration or melting of the accelerator agglomerates. During the drying operation the remaining water present in the product is removed, leaving an agglomerate of the powdered accelerator containing the rubber and water-insoluble solids content of the latex as the only diluents of the otherwise pure accelerator. If desired for purposes of identification, it is possible to produce colored agglomerated accelerators by the addition of small amounts of coloring agents to the aqueous slurry.

Further details of the practice of this invention are set forth in the following examples in which parts are shown by weight unless otherwise stated. These examples are to be interpreted as representative rather than restrictive of the score of this invention.

The 31.8 parts of the latex contained 10 parts of tribber hydrocarbon and 0.125 parts of styrenated pliciful as a non-discloring antioxidant. The diluted latex was added to an aqueous shirry of 2000 parts of water, 100 parts of 110 sodium chloride, I part of an alkylated aryl polyether alcohol and 190 parts of powdered mercapiobenzothazole. This combined mixture was flioroughly agitated and 100 piirts of 10% aqueous solution of alum 115 was added to coagulate the rubber latex. The excess water was decanted from the mixture and the residue washed 10 times with water, the filtered leaving a residue having a paste-like consistency. The paste like mixture was

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sodium alkyl aryl sulfonate as the emulsifier. The 45.2 parts of this latex contained 15.8 parts of rubber hydrocarbon and 0.2 parts of styrenated phenol as a non-discoloring antioxidant. This diluted latex was added to an aqueous slurry of 2000 parts of water, 100 parts of sodium chloride, 1.5 parts of dipolyoxyethylene alkyl tertiary amine, 100 parts of mercaptobenzothiazole and 200 parts of tetra-10 methyl thiuram disulfide. This combined mixture was thoroughly blended and the rubber latex coagulated with 100 parts of a 10% aqueous solution of alum. The excess water aqueous solution of alum. was decanted and the residue washed 10 times 15 with water, the excess water being decanted after each washing. After the last washing, the mixture was filtered leaving a residue of pastelike consistency which was extruded into string-like or rod-like shapes and dried at 20 125° F.

Example 3.

The same procedure was followed as in Example 1 except that the latex was made by the reaction of 55 parts of butadiene and 45 25 parts of acrylonitrile using a sodium alkyl aryl sulfonate as the emulsifier.

## EXAMPLE 4.

Water (350 parts) was added to 32.5 parts of a latex prepared from 75 parts of butadiene and 25 parts of styrene using 5 parts of a sodium alkyl aryl sulfonate as the emulsifier. The 32.5 parts of latex contained 11.35 parts of rubber hydrocarbon and 0.125 part of styrenated phenol. This diluted latex was added to an aqueous slurry of 1940 parts of water, 2720 parts of the sodium salt of mercaptobenzothiazole, and 90.8 parts of zinc sulfate. This combined mixture was thoroughly agitated and 150 parts of a 10% solution of sulfuric acid was added to coagulate the rubber latex. The excess water was decanted from the mixture and the residue repeatedly washed with water, the excess water being decanted after each washing. After the last washing the mixture was filtered, leaving a residue of pastelike consistency which was extruded into string-like or rod-like shapes and dried at 120° F.

Example 5. Water (375 parts) was added to 28.7 parts of a latex prepared from 75 parts of butadiene and 25 parts of styrene using 4 parts of a sodium alkyl aryl sulfonate as the emulsifier. The 28.7 parts of latex contained 10.2 parts of rubber hydrocarbon and 0.125 part of styrenated phenol. This diluted latex was added to an aqueous slurry of 2000 parts of water, 50 parts of sodium chloride and 190 parts of tetramethyl thiuram monosulfide. This combined mixture was thoroughly agitated and 10 parts of magnesium sulfate was added to coagulate the subber latex. The excess water was decanted from the mixture and the residue was washed repeatedly with water. After the

last washing the mixture was filtered, leaving a residue having a paste-like consistency which was extruded into rod-like shapes and dried at 125° F.

EXAMPLE 6.

Water (72.3 parts) was added to 28.7 parts of a latex prepared from 75 parts of butadiene and 25 parts of styrene using 4 parts of a sodium alkyl aryl sulfonate as the emulsifier. The 28.7 parts of latex contained 10.2 parts of rubber hydrocarbon and 0.125 part of styrenated phenol. This diluted latex was added to an aqueous slurry of 2000 parts of water, 50 parts of sodium chloride, and 190 parts of tetramethyl thiuram disulfide. This combined mixture was thoroughly agitated and 10 parts of alum was added to coagulate the rubber latex. The excess water was decanted from the mixture and the residue was repeatedly washed with water. After the last washing the mixture was filtered, leaving a residue having a paste-like consistency which was extruded into rod-like shapes and dried at 125° F.

The sodium alkyl aryl sulfonate employed in the examples is sold by National Aniline under

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the trade name Nacconol NR.

The agglomerated accelerators prepared according to the Examples 1 through 6 formed stable, non-dusting, rod-like agglomerates which were free-flowing. When these agglomerates were mixed with rubber on a mill or in a Banbury mixer, in accordance with normal practice, the agglomerates dispersed more rapidly and more uniformly than did the dry powdered accelerators from which agglomerates were made. Thus, it will be seen 100 that by following the practices of this invention, it is possible to produce a free-flowing, stable, non-dusting agglomerated accelerator containing only a minimum amount of diluent. The agglomerates themselves maintain their 105 stability during storage and handling operations and yet disperse rapidly and uniformly into and through the rubber compound with which they are used.

While certain representative embodiments 110 and details have been shown for the purpose of illustrating the invention, it will be apparent to those skilled in this art that various changes and modifications may be made therein without departing from the scope of the invention, as 115

defined in the appended claims.

What we claim is:-1. A method for preparing an agglomerated compounding ingredient for rubber characterized by the step of mixing an aqueous slurry of at least one powdered vulcanization accelerator selected from the group consisting of tetramethyl thiuram disulfide, tetraethyl thiuram, disulfide, tetramethyl thiuram dimethyl dithiocarmonosulfide, zinc bamate, zinc diethyl dithiocarbamate, zinc dibutyl dithiocarbamate, mercaptobenzothiazole, water insoluble metallic salts of mercaptobenzothiazole, and benzothiazyl disulfide with

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a latex stabilized against coagulation by at least 3 in which the powdered vulcanization accelerone acid-stable emulsifier as hereinbefore ator is benzothiazyl disulfide. defined and a rubber antioxidant, the latex 1511 5. A method according to any of Claims being selected from the group consisting of 1-3 in which the powdered vulcanization natural rubber latex, polychloroprene latex, polybutadiene latex, the latices of the rubbery copolymers of butadiene and styrene and the latices of the rubbery copolymers of butadiene and acrylonitrile, the mixture containing at 10 least 4% rubber hydrocarbon by weight of the accelerator, coagulating the rubber latex, removing the excess water, forming agglomerates of the mixture, drying the agglomerates to remove remaining water and collecting the 15 dried agglomerates.

2. A method according to Claim 1 in which - the acid-stable emulsifier is a synthetic organic material and the rubber antioxidant is non-discoloring.

3. A process according to Claim 1 or 2 in which the mixture is extruded to form strings or rods after the excess water is removed, drying the strings or rods to remove remaining water and collecting the dried material.

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25. 4. A method according to any of Claims 1-

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accelerator is mercaptobenzothiazole.

6. A method according to any of Claims 1-3 in which the powdered vulcanization accelerator is the zinc salt of mercaptobenzo-

7. A method according to any of Claims -6 in which a latex of the rubbery copolymer of butadiene and styrene is employed.

8. A method according to any of Claims 1-7 in which the latex stabilizer is a sodium

salt of an alkyl aryl sulfonate:
9. A method according to any preceding claim in which the amount of rubber or rubbery material in the latex does not exceed onetenth of the weight of the vulcanization accelerator, and is preferably in the range 4 6% by weight thereof.

10. A method for preparing an agglomerated compounding ingredient for rubber substantially as set forth and described hereinbefore.

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